## NEW UTILITY PATENT APPLICATION TRANSMITTAL (Large Entity)

(Only for new nonprovisional applications under 37 CFR 1.53(b))

Docket No. 1798-7267

Total Pages in this Submission

### TO THE ASSISTANT COMMISSIONER FOR PATENTS

**Box Patent Application** Washington, D.C. 20231

Transmitted herewith for filing under 35 U.S.C. 111(a) and 37 C.F.R. 1.53(b) is a new utility patent application for an

	ition e		ONS FOR DRYING SOLID SURFACES				
and ir	nvente	ed by	r,				
			AUD, 35 Boulevard Pasteur, 95210 Saint-Gratien, FRANCE				
es.							
lf a C	CONT	INUA	ATION APPLICATION, check appropriate box and supply the requisite information:				
	Conti	nuat	tion Divisional Continuation-in-part (CIP) of prior application No.:				
Enclo	osed a	аге:	Appellant W				
45			Application Elements				
1.	×	Filir	ng fee as calculated and transmitted as described below				
2.	×	Spe	edification having 18 pages and including the following:				
## ## #4	a.	×	Descriptive Title of the Invention				
11	b.		Cross References to Related Applications (if applicable)				
	c.		Statement Regarding Federally-sponsored Research/Development (if applicable)				
	d.		Reference to Microfiche Appendix (if applicable)				
	e.	×	Background of the Invention				
	f.	×	Brief Summary of the Invention				
	g.		Brief Description of the Drawings (if drawings filed)				
	h. 🛛 Detailed Description						
	i. 🛛 Claim(s) as Classified Below						
	j.	×	Abstract of the Disclosure				
3.		Drav	wing(s) (when necessary as prescribed by 35 USC 113)				
	a.		Formal				
	b.		Informal				
			Number of Sheets				

# NEW UTILITY PATENT APPLICATION TRANSMITTAL (Large Entity)

(Only for new nonprovisional applications under 37 CFR 1.53(b))

Docket No. 1798-7267

Total Pages in this Submission

### Application Elements (Continued)

4.	$\boxtimes$	Oath	or Declaration				
	a.		Newly executed (original or copy)   Unexecuted				
	b.		Copy from a prior application (37 CFR 1.63(d)) (for continuation/divisional application only)				
	C.	× ×	With Power of Attorney   Without Power of Attorney				
5.		The e	Incorporation By Reference (usable if Box 4b is checked) The entire disclosure of the prior application, from which a copy of the oath or declaration is supplied under Box 4b, is considered as being part of the disclosure of the accompanying application and is hereby incorporated by reference therein.				
6.		Com	outer Program in Microfiche ( <i>Appendix</i> )				
7.		Nucle	eotide and/or Amino Acid Sequence Submission (if applicable, all must be included)				
	a.		Paper Copy				
	b.		Computer Readable Copy (identical to computer copy)				
	c.		Statement Verifying Identical Paper and Computer Readable Copy				
			Accompanying Application Parts				
8.		Assig	gnment Papers (cover sheet & document(s))				
9.		37 C	FR 3.73(B) Statement (when there is an assignee)				
10.		Engli	ish Translation Document (if applicable)				
11.	×	Infon	mation Disclosure Statement/PTO-1449 🔻 Copies of IDS Citations				
12.	×	Preliminary Amendment					
13.	×	Acknowledgment postcard					
14.		Certi	ficate of Mailing				
			First Class   Express Mail (Specify Label No.):				
15.	×	Certi	ified Copy of Priority Document(s) (if foreign priority is claimed)				

# NEW UTILITY PATENT APPLICATION TRANSMITTAL (Large Entity)

(Only for new nonprovisional applications under 37 CFR 1.53(b))

Accompanying Application Parts (Cor

Docket No. 1798-7267

Total Pages in this Submission

Application Parts (Continued)	8. P
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16. Additio	onal Enclosures (p	lease identify be	low):		C542 U
		Fee Calcu	lation and Tra	nsmittal	
		CLAIMS	AS FILED		
For	#Filed	#Allowed	#Extra	Rate	Fee
otal Claims	15	-20 =	. 0	× \$22.00	\$0.00
dep. Claims	1	- 3 =	0	× \$82.00	\$0.00
ultiple Depende	nt Claims (check	if applicable)			\$0.00
				BASI	C FEE \$790.00
OTHER FEE (spe	ecify purpose)				\$0.00
2				TOTAL FILING	3 FEE \$790.00
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FIELD OF THE INVENTION

The invention relates to the field of drying and the subject of the invention is more particularly compositions for removing the water which superficially wets solid surfaces, in particular electronic materials (printed or integrated circuits), precision mechanical components, glass substrates and military, aerospatial, aeronautical or medical equipment.

\*\*BACKGROUND OF THE INVENTION\*\*

The usual process for dewetting solid

10 surfaces consists in immersing the wet substrate in an organic solution of an ionic or non-ionic surface-active agent. Until recently, the organic solvent commonly used was 1,1,2-trichloro-1,2,2-trifluoroethane (CFC 113) but its use has been banned because of its harmful effect on the stratospheric ozone laver.

The use of 1-chloro-1,1-difluoroethane
(HCFC 141b), provided as substitute for CFC 113 in
dewetting compositions, is already regulated and before
long it will no longer be possible to use this
20 compound.

To replace CFC 113 and HCFC 141b, the use has been envisaged of fluorinated hydrocarbons which, like 1,1,1,3,3-pentafluorobutane (365 mfc), 1,1,1,2,2,4,4-heptafluorobutane (347 mcf), (perfluorobutyl)ethylene (C<sub>4</sub>F<sub>9</sub>CH=CH<sub>2</sub>), 1H-perfluorohexane, n-perfluorohexane (PF 5060) or 1,1,1,2,3,4,4,5,5,5-decafluoropentane (43-10 mee), and perfluoro(methylmorpholine) (PF 5052), have no effect on the stratospheric ozone layer (zero

ODP), do not exhibit a flash point and, as is shown in the following table, possess physical characteristics which are similar to those of CFC 113 and HCFC 141b.

Solvent	Boiling point	Surface tension at 25°C (mN/m)	Relative density at 20°C
CFC 113	47.6	17.7	1.57
HCFC 141b	32	19.5	1.23
365 mfc	40	16.2	1.27
347 mcf	33.3	14.2	1.42
$C_4F_9CH=CH_2$	59	13.6	1.46
C <sub>6</sub> F <sub>13</sub> H	71	13	1.70
PF 5060	56	12	1.68
PF 5052	50	13	1.70
43-10 mee	54	14.1	1.58

However, the replacement of CFC 113 or of HCFC 141b by a fluorinated hydrocarbon such as those mentioned above is unsatisfactory because ionic-surface-active agents, which are most often used in dewetting compositions, are insoluble therein. Non-ionic surface-active agents are soluble in these fluorinated hydrocarbons but their dewetting effectiveness is often unsatisfactory.

To solve this problem in the case of 365 mfc, provision has been made, in the publication of Patent
15 JP 05-168805, for the addition of a cosolvent chosen

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from alcohols, ketones, alkanes and ethers. However, when the cosolvent is an alcohol comprising 1 to 3 carbon atoms, or a ketone, it is rapidly removed with the water, causing the rapid precipitation of the 5 surface-active agent. As regards alkanes and ethers, they exhibit a flash point, which presents a safety

problem for dewetting plants.  $_{NVEV}$  7000  $_{NE}$   $_{NVEV}$   $_{NVEV}$   $_{NEV}$   $_{NVEV}$   $_{NEV}$   $_{NEV}$ 

The subject of the present invention is thus, as composition useful for the dewetting of solid surfaces, a solution of at least one surface-active agent in a mixture of at least one fluorinated solvent and of at least one polyfluorinated alcohol of general formula:

 $Rf - (CH_2)_p - OH$  (I)

in which n is equal to 1 or 2 and Rf represents a linear or branched perfluoroalkyl radical containing from 4 to 8 carbon atoms. Such a composition does not exhibit a flash point under the standard determination conditions (ASTM standard D 3828).

In accordance with the present invention, the fluorinated solvent, which is liquid at room temperature, must not contain halogen atoms other than

fluorine but it can contain heteroatoms, such as oxygen or nitrogen. Use is advantageously made of a fluorinated solvent with a normal boiling point of between 20 and 100°C, preferably between 30 and 75°C.

5 Saturated or unsaturated fluorinated hydrocarbons containing from 3 to 6 carbon atoms, such as, for example, the compounds 365 mfc, 347 mcf, C<sub>4</sub>F<sub>9</sub>CH=CH<sub>2</sub>, PF 5060 and 43-10 mee mentioned above, as well as 1H-perfluorohexane (C<sub>6</sub>F<sub>13</sub>H) and 1,1,1,2,2,3,3,4,4-10 nonafluorohexane, are particularly well suited.

The polyfluorinated alcohols of formula (I) are known products. Among these, it is preferable to use those in which n is equal to 2. A more particularly preferred polyfluorinated alcohol is

15 tridecafluorooctanol ( $C_6F_{13}CH_2CH_2OH$  hereinafter TDFO).

The amount of polyfluorinated alcohol
necessary to completely dissolve the surface-active
agent in the dewetting composition according to the
invention can vary within wide limits, depending on the
20 nature of the fluorinated solvent and of the surfaceactive agent employed and according to the desired
concentration of surface-active agent. For a normal
concentration of surface-active agent (0.01 to 0.5% by
weight), a content of polyfluorinated alcohol ranging
25 up to 30% by weight is generally sufficient to produce
complete dissolution of the surface-active agent; in
certain cases, even a very low proportion of
polyfluorinated alcohol (0.1%) proves to be sufficient.

Consequently, in the dewetting composition according to the invention, the content of polyfluorinated alcohol can range from 0.1 to 30% by weight but it is advantageously between 0.5 and 5%.

- In the composition according to the invention, the surface-active agent can be any surface-active agent commonly used in dewetting compositions. Thus, non-ionic surface-active agents, such as fatty esters (for example, caprylamine caprylate and caprylamine, dodecylamine or stearylamine caproate), or fluorinated non-ionic surface-active agents, such as, for example, adducts of ethylene oxide with a polyfluorinated alcohol of formula (I), esters of polyfluorocarboxylic acids,
- 15 (polyfluoroalkyl) aminoalcohols, polyfluoroalkyl sulphides or polysulphides, or polyfluoroalkanecarbonamides or -sulphonamides, can be used. It is also possible to use anionic surface-active agents (for example a polyfluorosulphonic acid) but it
  20 is preferable to use cationic surface-active agents and more particularly those obtained by reaction of a monoor dialkyl phosphoric acid of general formula:

in which p is a number ranging from 1 to 2 and R

25 denotes a linear or branched alkyl radical containing
from 1 to 18 carbon atoms, with a quaternary ammonium
chloride of general formula:

$$R'_2N^{\dagger}R''_2Cl^{-}$$
 (III)

in which R' and R", which are identical or different, each represent a hydrogen atom or an alkyl or hydroxyalkyl radical containing 1 to 4 carbon atoms, and a fluorinated amine of general formula:

 $R_P - X - NR^1R^2$  (TV

in which R<sub>F</sub> represents a linear perfluoroalkyl radical containing from 2 to 20 carbon atoms, X represents a divalent bridge and the symbols R<sup>1</sup> and R<sup>2</sup>, which are identical or different, each represent a hydrogen atom 10 or an alkyl or hydroxyalkyl radical containing 1 to 4 carbon atoms.

Such cationic surface-active agents and their preparation are described in Patents FR 2,522,007 and 2,691,473, the contents of which are incorporated here by reference. As in the abovementioned patents, it is preferable to use:

- acids (II) in which R is a butyl, hexyl,
   2-ethylhexyl, octyl or tridecyl radical and more particularly mixtures of monoalkyl phosphoric acid and
   of dialkyl phosphoric acid in which the proportion by weight of the two acids is between approximately 60/40 and 40/60.
- quaternary ammonium chlorides (III) in
  which R' is a dodecyl or octadecyl radical and R" is a
  25 methyl radical or the commercial products mostly
  containing these chlorides,
  - fluorinated amines of formula:  $R_FC_2H_4SO_2NHCH_2CH_2CH_2N\left(CH_3\right){}_2$

#### RFC2H4CONHCH2CH2CH2N(CH3)2

As in the compositions of the prior art, the content of surface-active agent(s) in the dewetting

5 composition according to the invention can vary within wide limits. This content is generally low (from 0.01 to 0.5% by weight, preferably 0.04 to 0.2%) in readyfor-use dewetting compositions but it can reach up to 30% by weight in concentrated compositions which only 10 have to be diluted for the dewetting operation.

The solid surfaces from which water can be removed by means of a composition according to the invention can be highly varied, provided, however, that their substance (glass, metal, refractory material, plastic, and the like) is insoluble in and cannot be attacked by the fluorinated solvent(s) of the composition according to the invention.

EXAMPLES

The following examples illustrate the

The following examples illustrate the invention without limiting it. The cationic surface-active agents SA1 and SA2 were prepared from the following commercial products:

- butyl phosphoric acid  $(C_4H_9O)_{1.5}(OH)_{0.5}PO_2H$  exhibiting the following acid values:

strong acidity: 0.566 equivalent per 100 g

weak acidity: 0.276 equivalent per 100 g

which corresponds to a mixture containing approximately

40% of monobutyl phosphoric acid and 60% of dibutyl

phosphoric acid.

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0.4 g).

#### - Noramium M2C: mixture of different

homologues of formula:

R'2N+CH3)2 Cl

in which the distribution of the fatty chains (R') is

5	as follows:	C <sub>8</sub>	3%
		C <sub>10</sub>	6%
		C <sub>12</sub>	56%
		C <sub>14</sub>	18%
		C <sub>16</sub>	10%
10		C <sub>18</sub> , saturated	l 2%
		C18, unsaturat	ed 5%

This commercial product contains 75% of active material and contains 0.161 chloride equivalent per 100 g.

#### 15 Preparation of the surface-active agent SA1

2.69 g of butyl phosphoric acid, 3.91 g (i.e.

- 7.6 meq.) of the fluorinated amine  $C_6F_{13}C_2H_4SO_2NHCH_2CH_2CH_2CH_2$  (CH<sub>3</sub>)<sub>2</sub>, 3 g of a 10% methanolic sodium hydroxide solution (i.e. 7.6 meq. NaOH) and 8.32 g of Noramium 20 M2C are added successively and with stirring to 150 g of HCFC 141b. This final addition causes the immediate formation of a sodium chloride precipitate which is removed by filtration (weight of NaCl collected:
- 25 After removing the sodium chloride precipitate, the filtrate is then evaporated to dryness at 110°C in an oven. 14.8 g of surface-active agent SA1 are thus obtained.

### Preparation of the surface-active agent SA2

The preparation is carried out as above but replacing the 150 g of HCFC 141b with 300 g of TDFO and dispensing with the final stage of evaporation to dryness. A solution containing approximately 5% of surface-active agent SA2 in TDFO is thus obtained.

#### EXAMPLE 1

A dewetting solution is prepared by dissolving 100 mg of surface-active agent SA1 in a 10 mixture of 95 g of HFC 365 mfc and 5 g of TDFO and then this solution is introduced into a beaker placed in a small Branson 2200 ultrasonic vessel.

A 20  $\times$  50 mm stainless steel grid, wetted beforehand by immersion in water, is then immersed in the dewetting solution and maintained under ultrasonic agitation for one minute.

After having been withdrawn from the dewetting solution, the grid is immersed in absolute alcohol and the quantitative determination of water by 20 the Karl Fischer method indicates that 94% of the water has been removed.

The same result is obtained if the fluorinated alcohol TDFO is replaced with the same amount of a technical mixture of alcohols of formula  $25 \quad Rf-CH_2CH_2-OH, \ \, \text{the distribution of the Rf radicals being centred about } C_6.$ 

If the fluorinated alcohol TDFO is replaced with the same amount of n-pentanol or of n-hexanol, the

degree of removal of the water is only 60% and 72% respectively.

#### EXAMPLE 2

A dewetting solution is prepared by mixing

5 1 g of the solution containing 5% of surface-active
agent SA2 and 99 g of HFC 365 mfc. A dewetting solution
containing 500 ppm of surface-active agent SA2 is thus
obtained, which solution is used as in Example 1. The
degree of removal of the water is 95%.

Substantially the same result is obtained by using a dewetting solution containing 0.1% of surface-active agent SA2 in a mixture of 99 g of HFC 365 mfc and 1 g of TDFO.

#### EXAMPLES 3 to 11

The procedure is as in Example 2, but replacing the solvent HFC 365 mfc with other fluorinated solvents. The operating conditions and the results are combined in the following table.

Example	Solvent	(%)	% of TDFO	SA2 (ppm)	Water removed
3	347 mcf	98	2	500	93
4	C <sub>4</sub> F <sub>9</sub> CH=CH <sub>2</sub>	95	5	1000	92
5	C <sub>4</sub> F <sub>9</sub> CH=CH <sub>2</sub>	99	1	500	95
6	C4F9CH2CH3	99	1	500	91
7	C <sub>4</sub> F <sub>9</sub> CH <sub>2</sub> CH <sub>3</sub>	98	2	500	92
8	C <sub>6</sub> F <sub>13</sub> H	98	2	500	91
9	PF 5060	99	1	500	92
10	PF 5052	99	1	500	94
11	43-10 mee	98	2	500	94

#### EXAMPLE 12

The procedure is as in Example 1, but replacing the fluorinated alcohol TDFO with

5 pentafluoropropanol (CF<sub>3</sub>CF<sub>2</sub>CH<sub>2</sub>OH, hereinafter PFP) and using a dewetting solution obtained by dissolving

100 mg of surface-active agent SA1 in a mixture of 80 g of HFC 365 mfc and 20 g of PFP. The degree of removal of the water is 95%.

#### EXAMPLE 13

The procedure is as in Example 1, but using a dewetting solution obtained by dissolving 100 mg of N,N'-dioleoyl-N-oleylpropylenediamine (hereinafter surfactant SA3) in a mixture of 90 g of HFC 365 mfc and

10 g of TDFO. The degree of removal of the water is 91% .

If the TDFO is replaced with the same amount  $(10\ g)$  of PFP, the degree of removal of the water is  $5\ 93\%$ .

#### EXAMPLE 14

The procedure is as in Example 1, but using a dewetting solution obtained by dissolving 100 mg of the adduct of 6 mol of ethylene oxide with one mole of TDFO in a mixture of 99 g of HFC 365 mfc and 1 g of TDFO.

The degree of removal of the water is 90%.

If the mixture of 365 mfc and of TDFO is replaced with a mixture of 90 g of 365 mfc and 10 g of PFP, the degree of removal of the water is also 90%.

#### 15 **EXAMPLES 15 to 18**

The procedure is as in Example 1, but replacing the fluorinated alcohol TDFO with PFP and using a dewetting solution obtained by dissolving, in a mixture of 80 g of HFC 365 mfc and 20 g of fluorinated 20 alcohol PFP, 100 mg of one of the following surfaceactive agents:

 $SA4 = C_6F_{13}C_2H_4(OC_2H_4)_6OH$ 

 $SA5 = C_8F_{17}C_2H_4N(C_2H_4OH)_2$ 

25 SA6 =  $C_6F_{13}C_2H_4SO_2NHCH_2CH_2CH_2N(CH_3)_2$ 

 $SA7 = C_6F_{13}C_2H_4SO_3H$ ,  $H_2NCH_2CH(C_2H_5)C_4H_9$ 

The results obtained are combined in the following table.

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Example	Surface-active agent	Water removed (%)					
15	SA4	93					
16	SA5	93					
17	SA6	93					
18	SA7	95					

#### EXAMPLE 19

22 litres of the dewetting solution of
Example 2 are charged to the dewetting and settling
tanks of a drying machine with a capacity of 33 litres
5 and 11 litres of pure HFC 365 mfc are charged to the
rinsing vessel.

A wet stainless steel grid, resulting from a prior washing and rinsing treatment in aqueous medium, is immersed in the dewetting tank for 3 minutes and then rinsed for 3 minutes in the pure HFC 365 mfc at boiling point.

The grid is subsequently immersed in 100 g of

15 absolute ethanol, the water content of which was
determined beforehand by the Karl Fischer method. A new
quantitative determination carried out after immersion
of the grid revealed no increase in the water content,
which means that the composition according to the

20 invention makes it possible to obtain virtually
complete removal of the water.

Although the invention has been described in conjunction with specific embodiments, it is evident that many alternatives and variations will be apparent to those skilled in the art in light of the foregoing description. Accordingly, the invention is intended to embrace all of the alternatives and variations that fall within the spirit and scope of the appended claims. The above references are hereby incorporated by

10 reference.

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Sub Pl Dewetting composition, characterized in that it is composed of a solution of at least one surface-active agent in a mixture of at least one.
 fluorinated solvent and of at least one polyfluorinated alcohol of general formula:

$$Rf-(CH_2)$$
 OH (I)

in which n is equal to 1 or 2 and Rf represents a linear or branched perfluoroalkyl radical containing

- 10 from 4 to 8 carbon atoms.
  - Composition according to Claim 1,
    containing at least one alcohol of formula (I) in which
    n is equal to 2.

3. Composition according to Claim 1, in. which the alcohol of formula (I) is tridecafluorooctanol (C<sub>6</sub>F<sub>12</sub>CH<sub>2</sub>CH<sub>2</sub>OH).

5 who

- 4. Composition according to one of Claims 1 to 3, in which the fluorinated solvent has a normal boiling point of between 20 and 100°C, preferably
- 0 between 30 and 75°C.

355 1 P

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5. Composition according to claim 4, inwhich the fluorinated solvent is a saturated or unsaturated fluorinated hydrocarbon containing from 3 to 6 carbon atoms.

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6. Composition according to Claim 4 or 5, in which the fluorinated hydrocarbon is chosen from 1,1,1,3,3-pentafluorobutane, 1,1,1,2,3,4,4,5,5,5-

decafluoropentane, 1,1,1,2,2,3,3,4,4-nonafluorohexane, 1H-perfluorohexane, n-perfluorohexane, (perfluorobutyl)ethylene and perfluoro (methylmorpholine).

7. Composition according to one of Claims 1 to 6, in which the surface-active agent is a cationic surface-active agent obtained by reaction of a mono- or dialkyl phosphoric acid of general formula:

10 in which p is a number ranging from 1 to 2 and R denotes a linear or branched alkyl radical containing from 1 to 18 carbon atoms, with a quaternary ammonium chloride of general formula:

$$R'_2N^*R''_2C1^-$$
 (III)

in which R' and R", which are identical or different, each represent a hydrogen atom or an alkyl or hydroxyalkyl radical containing 1 to 4 carbon atoms, and a fluorinated amine of general formula:

$$R_F - X - NR^1R^2$$
 (IV)

- 20 in which R, represents a linear perfluoroalkyl radical containing from 2 to 20 carbon atoms, X represents a divalent bridge and the symbols R<sup>1</sup> and R<sup>2</sup>, which are identical or different, each represent a hydrogen atom or an alkyl or hydroxyalkyl radical containing 1 to
- 25 4 carbon atoms.
  - 8. Composition according to Claim 7, in which R is a butyl, hexyl, 2-ethylhexyl, octyl or tridecyl radical R' is a dodecyl or octadecyl radical,

Syn A

R" is a methyl radical. X is a -CH<sub>2</sub>CH<sub>2</sub>SO<sub>2</sub>NHCH<sub>2</sub>CH<sub>2</sub>- or -C<sub>2</sub>H<sub>4</sub>CONHCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>- bridge and R<sup>1</sup> and R<sup>2</sup> are methyl radicals.

- 9. Composition according to one of Claims 1
  5 to 8, in which the content of polyfluorinated
  alcohol(s) is between 0.1 and 30% by weight, preferably
  between 0.5 and 5%
  - 10. Composition according to one of Claims 1 to 9, in which the content of surface-active agent(s) is between 0.01 and 0.5% by weight, preferably between 0.04 and 0.2%.
  - 11. Composition according to one of Claims 1 to 9, in the form of a concentrate containing up to 30% by weight of surface-active agent(s).
  - 12. Use of a composition according to one of Claims 1 to 10 for the dewetting of solid surfaces.

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ROGO

PATENT APPLICATION

COUNTRY:

FRANCE

Filed on:

4 March 1997

No.:

97.02550

File No.:

AM 1163

#### TEXT FOR FOREIGN FILING

#### ABSTRACT

#### 0000000

#### COMPOSITIONS FOR DRYING SOLID SURFACES

#### 0000000

Company called:

ELF ATOCHEM S.A.

4/8 Cours Michelet

La Défense 10

92800 PUTEAUX

#### 0000000

Agent:

Jean LEBOULENGER

0000000

Inventor:

Pascal MICHAUD

#### 0000000

A drying composition according to the invention is composed of a solution of a surface-active agent in a mixture of a fluorinated hydrocarbon and of a polyfluorinated alcohol of formula:

$$Rf - (CH_2)_n - OH$$
 (I)

in which n is equal to 1 or 2 and Rf represents a linear perfluoroalkyl radical containing from 2 to 8 carbon atoms.



Docket No. 1798-7267

[ ] YES [ ] NO

# COMBINED DECLARATION AND POWER OF ATTORNEY FOR ORIGINAL, DESIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL, DIVISIONAL, CONTINUATION OR CONTINUATION-IN-PART APPLICATION

As a below named inventor, I hereby declare that:

COMPOSITIONS FOR DRYING SOLID SURFACES

the specification of which

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

	a. [X]	is attached here	eto				
	b. [ ]			as application Serial No	o and e).		
	PCT FILED APPLICATION ENTERING NATIONAL STAGE						
	c. [ ]			tional Application No.			
			ed and understand the mendment referred to		dentified specification, including		
			ose information which of Federal Regulations		nation of this application in		
-	I hereby claim foreign priority benefits under Title 35, United States Code § 119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:						
	The attached 35 U.S.C. § 119 claim for priority for the U.S. application(s) listed below forms a part of this declaration.						
Country		Application Number	Date of filing (day, month, yr)	Date of issue (day, month, yr)	Priority Claimed		
France		97.02550	04 March 1997		[x] YES [ ] NO		
					[ ] YES [ ] NO		

Docket No. 1798-7267

# ADDITIONAL STATEMENTS FOR DIVISIONAL, CONTINUATION OR CONTINUATION-IN-PART

I hereby claim the benefit under Title 35, United States Code § 120 of any United States application(s) listed below.

Application Serial No.	Filing Date	Status (patented, pending, abandoned)
Application Serial No.	Filing Date	Status (patented, pending, abandoned)

[] In this continuation-in-part application, insofar as the subject matter of any of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, § 1.56(a) which occurred between the filling date of the prior application and the national or PCT international filling date of this application.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or Imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

I hereby appoint the following attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewiti: Frederick F. Calvetti (28,557), BELL, BOYD & LLOYD, P.O. Box 65331, Washington, DC 20035-5331, Telephone: (202) 466-6300, Telefax: (202) 463-0678. Address all correspondence to BELL, BOYD & LLOYD, P.O. Box 65331, Washington, D.C. 20035-5331.

The undersigned hereby authorizes the U.S. attorneys named herein to accept and follow instructions from the undersigned's assignee, if any, and/or, if the undersigned is not a resident of the United States, the undersigned's domestic attorney, patent attorney or patent agent, as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorneys and the undersigned. In the event of a change in the person(s) from whom instructions may be taken, the U.S. attorneys named herein will be so notified by the undersigned.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under \$1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

I hereby specify the following as the correspondence address to which all communications about this application are to be directed:

Post Office Address \_\_\_\_

	SEND CORRESPONDENCE TO:	
	BELL, BOYD & LLOYD P.O. BOX 65331 Washington, DC 20035-5331.	
	DIRECT TELEPHONE CALLS TO: Frederick F. Calvetti, Tel: 20	02-466-6300
Inver Resid	name of sole or first inventor Pascal MICHAUD  ntor's signature*  lence Saint-Gratien, FRANCE  enship French  Office Address 35 Boulevard Pasteur, 95210 Saint-Gratien, France	March, 3, 1998 date
Full	name of second joint inventor, if any	
Resid	ntor's signature*enship	date